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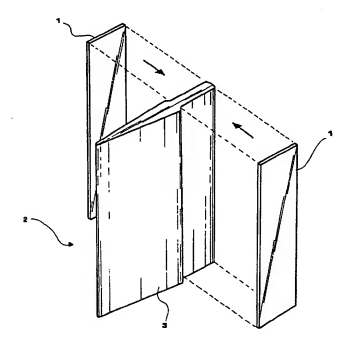
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(54) Process for the manufacturing of ceramic-matrix composite layers and related composite material

(57) Process for the manufacturing of ceramic-matrix composite layers resistant to very high temperatures, comprising the steps of: preparing the powders for the feeding of the deposition plant by wet mixing of the ceramics constituting the material in form of fine powders and atomizing the suspension in the presence

of a hot air jet; and depositing by plasma thermal spraying with an inert gas flow and with a > 30 kPa pressure in a deposition chamber. This process forms composite layers having a very high resistance, apt to be employed as coatings for vehicles, of the type destined to reenter Earth atmosphere from outer space and to be launched again therein.



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Description

[0001] The present invention refers to a process for the manufacturing of ceramic-matrix composite layers resistant to very high temperatures, like e.g. Boride mixtures, like e.g. Zirconium, Titanium and/or Hafnium Borides, or Oxide mixtures, like e.g. Hafnium and/or Zirconium Oxides, and Carbides, like e.g. Silicon, Tantalum, Zirconium and/or Hafnium Carbides. These layers can serve as protective coating of the material onto which they are coated, or they can be employed as components per se, upon having removed the substrate material, e.g. by machining or by chemical attack. Hence, the present invention further refers to the composite material obtained by said process.

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[0002] Of late years, a specific class of materials, that of the Ultra High Temperature Ceramics (UHTC) has aroused a marked interest for prospective employs thereof in the aerospace field.

[0003] In particular, a group of composites having as matrix the Borides of some transition metals (ZrB₂ and HfB₂) is deemed to be the most suitable to the thermal protection of sharp leading edges for the new generations of space vehicles, apt to reenter Earth atmosphere after a stay thereof in outer space and to be launched again therein.

[0004] The context of said viable and exemplary employ of the materials subject-matter of the present invention will be detailed hereinafter.

[0005] The material specifications required by such a specific employ are extremely complex and difficult to attain all in the same material. In fact, the materials at issue should combine high thermal conductivity coefficients and high resistance to thermal shocks to a very high melting point and a low thermal expansion coefficient..

The high melting point, apt to reach the 3300K, is a real obstacle to the shaping of these materials. In fact, the sole technique capable of ensuring satisfactory features in terms of porosity and of mechanical resistance is the hot pressing, entailing, however, remarkable limitations in terms of sizes and costs.

[0006] Other shaping techniques are known. E.g., the plasma thermal spraying overcomes the problems entailed in the hot pressing forming techniques of the state of the art. In fact, the employ of a plasma that in its hotter region could reach and exceed the 20000K enables the deposition of virtually all those elements or compounds having a stable liquid phase. However, it should be borne in mind that the manufacturing of self-standing, high-melting ceramics-based coatings and components still constitutes, due to the required powers, a frontier research field in the employ of the plasma thermal spraying. Moreover, depositing a composite material entails the risk of interactions between the constituent phases thereof.

[0007] The present invention consists of a methodology of depositing by plasma thermal spraying, yielding

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Boron (Boride)-based, Oxygen (Oxide)-based, Boride and Oxide mixtures-based, and Carbon (Carbide)-based ceramic-matrix composite materials. These composites are characterized by the concomitant presence of highly compact ceramics phases (density >96% of the theoretical density).

[0008] The technical problem underlying the present invention is to attain these features, overcoming the obstacles insurmountable by the state of the art.

[0009] This problem is solved by a process as abovespecified, comprising the steps of:

- preparing the powders for the feeding of the deposition plant by wet mixing the ceramics constituting the material in form of fine powders and atomizing the suspension in the presence of a hot air jet;
- depositing by plasma thermal spraying with an inert gas flow and with a >30 kPa pressure in a deposition chamber.

[0010] The main advantage of the process according to the invention lies in forming complex-shaped and self-standing composite layers suitable to the most extreme employs, as in the coatings of space vehicles.

[0011] The present invention will hereinafter be described according to preferred embodiments thereof, given as a non-limiting example with reference to examples reported hereinafter and to the individual Figure attached, which depicts a component coated according to the abovedefined process.

[0012] The process subject-matter of the invention is based on plasma spraying, in which a plasma generated by an electric arc is employed to melt and to accelerate powders of the materials to be deposited. Hence, the process comprises the methodology for preparing the starting powders and the selection of the spraying process parameters.

[0013] The powders are prepared mixing thereamong the different ceramics materials, in form of powders having a <10 μ m average particle size so as to attain powders made of particles having a >10 μ m average diameter and a satisfactory flowability.

[0014] The different ceramics materials are destined to form the different phases of a ceramic-matrix composite layer.

[0015] The powders are made of two distinct ceramics materials, the first one being Boron (Boride)-based, Oxygen (Oxides)-based, and based on mixtures thereof, and the second one being Carbon (Carbide)-based.

[0016] The first material may be selected from a group of compounds comprising Zirconium Boride, Titanium Boride, Hafnium Boride, Zirconium Oxide, Hafnium Oxide, and mixtures thereof.

[0017] The second material may be selected from a group of compounds comprising Silicon Carbide, Titanium Carbide, Tantalum Carbide, Zirconium Carbide, Hafnium Carbide, and mixtures thereof.

[0018] The ratio among these materials may vary ac-

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cording to the needs and to the properties to be conferred to the ceramic matrix. In the present embodiment, the first material is present in a 40% to 95% ratio (preferred: about 75%), and the second material is present in a 5% to 60% ratio (preferred: about 25%). It is understood that said ratios should be compatible with the structure provided for the ceramic matrix.

[0019] Said mixing may be wet, i.e., carried out, e.g., in an aqueous environment or in the presence of water-dissolved organic binders, e.g. of a polyvinyl binder.

[0020] The suspension resulting from said mixing may be dried and atomized, i.e. finely dispersed, nebulizing it with a >200kPa compressed hot-air jet, in particular to a >373K temperature, thereby obtaining a powder apt to be employed in a plasma spraying plant.

[0021] Said powders, prior to the plasma spraying, may be subjected to an optional thermal treatment fostering the sintering thereof.

[0022] The deposition of the layers of composite material, starting from the powders thus prepared, should take place under the following conditions:

- (a) in the presence of an inert, i.e. non-oxidizing, e. g., Argon (Ar), atmosphere in order to prevent particle reactions during the deposition; and
- (b) with a high-temperature and high-density plasma, in order to allow the complete melting of the particles.

[0023] The specific parameters vary according to the selected materials.

[0024] The deposition will take place under a non-oxidizing gas atmosphere, at a power such as to attain a plasma temperature sufficiently high to form a composite matrix. Parameters like the distance between the torch and the substrate to be coated, the powder feed flow rate, the number of required passes and the temperature to be conferred to the substrate will suitably be selected.

[0025] The pressure in the deposition chamber is >30kPa, and it preferably ranges from 120 to 300 kPa.
[0026] The plasma is generated by an electric arc having a >30kW power, and preferably ranging from 35 to 55 kW.

[0027] The employed plasma is formed lonizing a flow of Argon and Hydrogen mixtures, having a Argon flow rate of >20 SLPM (Standard Liters Per Minute), and preferably ranging from 35 to 55 SLPM, and having a Hydrogen flow rate of >6 SLPM, and preferably ranging from 8 to 15 SLPM.

[0028] Alternatively, the employed plasma is formed ionizing a flow of Argon and Helium mixtures, having an Argon flow rate of >10 SLPM, and preferably ranging from 15 to 30 SLPM, and having a Helium flow rate of >30 SLPM, and preferably ranging from 50 to 100 SLPM.

[0029] The coated substrate, post-deposition and after a suitable cooling thereof, may be subjected to op-

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tional machine finishing.

EXAMPLE 1

[0030] The deposition of a coating of UHTC material, consisting of a ZrB₂ matrix incorporating evenly distributed SiC particles, was carried out.

[0031] The process for the attainment thereof was based on the following steps: (i) ZrB_2 powders, having a 5 μ m average particle size, and SiC, having a 0.7 μ m average particle size, were mixed in an aqueous suspension, in the presence of a polyvinyl binder, in the ratios: 75 % (b/w) ZrB_2 and 25% SiC. Then, the suspension was atomized with a 520K compressed air flow;

(ii) the powders were sprayed with a ≤80 kW power plasma torch. Said torch was installed in a vacuum chamber, so as to control the atmosphere composition and pressure;

(iii)as substrate, a 40mm-diameter graphite tubular component was employed;

(iv) prior of starting the deposition, the chamber was scavenged up to a

2Pa vacuum level. Subsequently, Ar was inletted, up to a 200KPa pressure;

(v) then the deposition process was started, employing the following parameters:

- plasma generating gas flow: 47 SLPM Ar + 10 SLPM H₂ (SLPM = standard liters per minute).
- electric arc power: 42 kW.
- Torch -substrate distance: 110 mm.
- * powder feed flow rate: 7 g/min.
- * number of torch passes onto substrate: 75.
- * during the deposition the substrate did not surpass the temperature of 500K; this situation was attained cooling the substrate with an Ar flow at ambient temperature.
- 40 [0032] At the end of the deposition process the coated sample was extracted from the deposition chamber and subsequently tested with a thermal cycling test under inert atmosphere of between 600K and 2300K. Several cycles were carried out, in each thereof the component having been heated up to 2300K over a time equal to 200 s, held thereat for a time equal to 600 s, and finally cooled down to 600K over a time equal to 600 s.
- [0033] At the end of the test, there were carried out non-disruptive tests, and subsequently optical microscopy analyses, which did not highlight mechanical damage (like cricks and/or detachments from the substrate).

EXAMPLE 2

[0034] The deposition of a coating of UHTC material, consisting of a HfB₂ matrix incorporating evenly distributed SiC particles, was carried out.

[0035] The process for the attainment thereof was

based on the following steps:

(vi) HfB $_2$ powders, having a 5 μ m average particle size, and SiC, having a 0.7 μ m average particle size, were mixed in an aqueous suspension, in the presence of a polyvinyl binder, in the ratios: 75 % (b/w) ZrB $_2$ and 25% SiC. Then the suspension was atomized with a 520K compressed air flow;

(vii)The powders were sprayed with an ≤80 kW power plasma torch. Said torch was installed in a vacuum chamber, so as to control the atmospheric composition and pressure;

(viii) as substrate, a stainless steel AISI 416 sheet was employed;

(ix) prior of starting the deposition, the chamber was scavenged up to a 1 Pa vacuum level. Subsequently, Ar was inletted, up to a 220 kPa pressure;

(x) then the deposition process was started, employing the following parameters:

plasma generating gas flow: 40 SLPM Ar + 12
 SLPM H2 (SLPM -

Standard Liters Per Minute).

- * electric arc power: 45 kW.
- torch substrate distance: 110 mm.
- * powder feed flow rate: 10 g/min.
- * number of torch passes onto substrate: 80.
- during the deposition the substrate did not surpass the temperature of

400K; this situation was attained cooling the substrate with an Ar flow at ambient temperature.

[0036] At the end of the deposition process, the coated component was extracted from the deposition chamber. Then the substrate was removed by machining (milling), thereby obtaining a 1.2 mm thick HfB₂-SiC composite plate.

[0037] At one end of said plate, a bevel was obtained by diamond tool machining, so as to simulate a sharp leading edge.

[0038] In the Figure, a component 1 made with a coating 2 obtained with the abovedescribed process, with a characteristic sharp angle shape whose function will be detalled hereinafter, is sketched.

[0039] The component, apt to form the leading edge of a wing of an aerospace vehicle apt to repeatedly reenter Earth atmosphere, comprises suitable fastening means 3 to connect it to the wing structure.

[0040] Concerning the application of like coatings, the employ thereof in the aerospace industry is of the utmost interest.

[0041] In fact, the Earth atmosphere reentering stage, at the end of an orbital or interplanetary flight, is particularly severe for the vehicles, subjected to an intense thermal flow due to the resistance of the atmospheric strata. To date, for manned missions two reentry modes are provided: ballistic, usually not implying the reuse of

the vehicle, or airlift-supported.

[0042] A airlift-supported reentry implies that the vehicle have an adequate configuration, developing a non-nil lift/drag ratio (L/D>O); the adopted trajectory is long, and it enables, as long as maneuverability during the final stage of the flight is ensured, a conventional-type landing, with no vehicle damage compromising the reuse thereof.

[0043] By virtue of the longer stay in the atmosphere, the variation of kinetic energy is distributed over a lengthier time interval, thereby reducing the heat flow to which the structure is subjected. In this case, the thermal protections consist of thermally insulating shields operating under non-ablative conditions, which shield the inside of the vehicle and irradiate towards the outside thereof a substantial fraction of the absorbed heat. The application of ablative shields would prove inappropriate, as the materials available to date would not reach a temperature high enough to be up to their task, merely undergoing degradation.

[0044] The selection of the airlift-supported reentry has paved the way to the first generation of Reusable Launch Vehicles (RLV's), the Space Shuttles, since post-landing each Shuttle is nearly unscathed and it can subsequently be reused in further workcycles. However, aircraft operativeness is not free from problems, some of which being of pressing interest to the future development of Space activities, concerning the correlation between the thermal protection systems and the aerodynamic efficiency of the aircrafts.

[0045] A particularly relevant problem, affecting the operativeness of the first generation of RLV's lies in the modest aerodynamic efficiency, expressed by the Lift/ Drag ratio, of the edges adopted. Due to the blunt configuration of the nose and of the leading edges of the wings, the vehicles meet a remarkable drag which has to be compensated, during the launch, by a power surplus, and which entails a payload reduction.

[0046] Moreover, the selection of the reentry trajectory is complicated. However, to date this configuration is unavoidable in order to dispose of the heat generated by atmospheric friction. Hence, so far a different configuration, with sharp leading edges and pointed nose, has not proved viable, relevant advantages thereof notwithstanding.

[0047] Among said advantages, e.g., an Increased aerodynamic efficiency and therefore an increased payload, greater freedom in selecting the reentry orbits, lesser interferences due to the free electrons associated to the rounded edges.

[0048] In particular, the increased aerodynamic efficiency is due to the decreased contribution to the drag caused by the reduction of the related momentum associated to the fluid crossed.

[0049] The reason behind the discarding of the sharp configuration in hypersonic applications essentially lies in the irreconciliability between the aerothermal constraint imposed thereby and the performances of the

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available thermal protections. The latter generally attain ablative conditions, with the entailed rounding off of their edges and the decay of the efficiency features thereof. [0050] Therefore, in the X-15 rocket vehicles and in the Apollo and Shuttle missions the blunt configuration has mandatorily been adopted, it being less effective yet capable of protecting the structures from excessive thermal loads. By virtue of the detached shock system associated thereto, the heat flow generated by the change in momentum inside of the fluid does not directly burden the sharp leading edge surrounding it, thereby preventing the overheating thereof.

[0051] The above remarks highlight how some of the problems related to the reentry operativeness of the RLV's be closely linked to the performances of the materials currently made available by the aerospace technologles. Apparently, as the length of the ground inspection times, as well as the aerodynamic efficiency, via the configuration of the edges, depend thereupon, and as these issues are crucial to the future development of the Space activities, the quest for materials exhibiting improved performances is fundamental.

[0052] The composite material manufactured with the abovedescribed process, in particular from the ZrB₂-SiC and the HfB₂-SiC compounds, by virtue of the marked resistance thereof to high temperatures, can advantageously be employed in this context as composite coating layer.

[0053] The employ is made viable by the fact that they be suitable to the thermal protection of the sharp leading edges, thereby providing the implementation thereof in the future generations of RLV's.

[0054] To the abovedescribed process for the manufacturing and related material a person skilled in the art, in order to satisfy further and contingent needs, may effect several further modifications and variants, all however comprised in the protective scope of the present invention, as defined by the appended claims.

Claims

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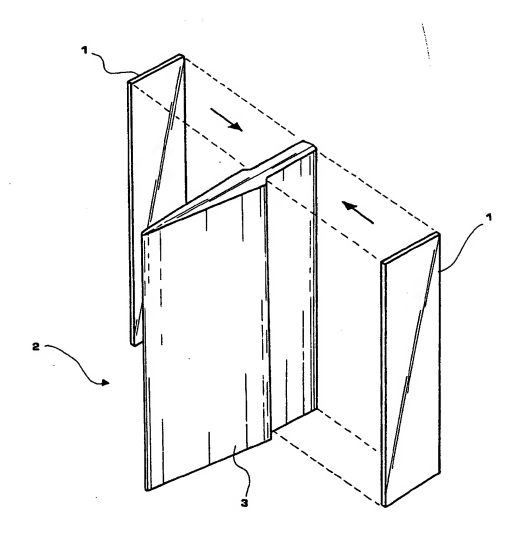
- A process for the manufacturing of ceramic-matrix composite layers resistant to very high temperatures, comprising the steps of:
 - preparing the powders for the feeding of the deposition plant by wet mixing of the ceramics constituting the material in form of fine powders and atomizing the suspension in the presence of a hot air jet;
 - depositing by plasma thermal spraying in a deposition chamber containing inert gas at a >30kPa pressure.
- The process according to claim 1, wherein the powders are prepared mixing thereamong the different ceramics materials, in form of powders having a > 10

 μm average particle size, so as to attain powders made of particles having a >10 μm average diameter

- 3. The process according to claim 1, wherein the powders are made of two distinct ceramics materials, the first material selected from a group of Boron (Boride)-based, Oxygen (Oxides)-based compounds, and mixtures thereof, and the second material being Carbon (carbide)-based.
- 4. The process according to claim 3, wherein the first material may be selected from a group of compounds comprising Zirconium Boride, Titanium Boride, Hafnium Boride, Zirconium Oxide, Hafnium Oxide, and mixtures thereof.
 - 5. The process according to claim 3, wherein the second material may be selected from a group of compounds comprising Silicon Carbide, Tantalium Carbide, Zirconium Carbide, Titanium Carbide, Hafnium Carbide, and mixtures thereof.
- 6. The process according to claim 3, wherein the first material is present in a 40% to 95% ratio, and the second material is present in a 5% to 60% ratio.
- The process according to claim 1, wherein said mixing is carried out in an aqueous environment and/ or in the presence of water-dissolved organic binders.
- The process according to claim 1, wherein said compressed hot air jet has a > 373K temperature.
- The process according to claim 1, wherein said inert gas contained in said deposition chamber comprises Argon, Helium.
- 40 10. The process according to claim 1, wherein said pressure in said deposition chamber preferably ranges from 120 to 300 kPa.
- 11. The process according to claim 1, wherein said plasma is generated by an electric arc having a >30 kW power, and preferably ranging from 35 to 55 kW.
 - 12. The process according to claim 1, wherein said plasma is formed ionizing a flow of Argon and Hydrogen mixtures, having a Argon flow rate of >20 SLPM, and preferably ranging from 35 to 55 SLPM, and having a Hydrogen flow rate of >6 SLPM, and preferably ranging from 8 to 15 SLPM.
- 5 13. The process according to claim 1, wherein said plasma is formed ionizing a flow of Argon and Helium mixtures, having a Argon flow rate of >10 SLPM, and preferably ranging from 15 to 30 SLPM,

and having a Helium flow rate of >30 SLPM, and preferably ranging from 50 to 100 SLPM.

- 14. The process according to claim 1, wherein, during the thermal spraying, the plasma has high temperature and density, such as to enable the complete sintering of the sprayed particles.
- **15.** A composite material manufactured with the process as defined by Claim 1.
- **16.** A coating, in particular for aerospace employs on reusable vehicles, comprising the composite material of the preceding claim.





EUROPEAN SEARCH REPORT

Application Number

EP 02 42 5143

A	US 4 645 716 A (HARRINGTON JOHN H ET 24 February 1987 (1987-02-24) - claims 7,15-18 * - examples 1,4 * US 6 110 853 A (BERGER LUTZ-MICHAEL ET LUTZ-MICHAEL E	3,4,6 T 1,2,7-16	C23C4/10 C04B35/58 C04B35/488 C04B41/87
A #	L) 29 August 2000 (2000-08-29) claims 19,25,26 *	-,-,. 20	
- 1	example 1 *	3,5,6	
2	S 3 617 358 A (DITTRICH FERDINAND J) November 1971 (1971-11-02) abstract * column 6, line 22 - line 30 *	1,3,7,8,	
1:	E 197 14 433 A (HOECHST AG) 5 October 1998 (1998-10-15) claims 1,4,8,17,18 * examples *	1-4,6, 9-16	TECHNICAL FIELDS SEARCHED (Int.Cl.7)
DE SI CE PR vo 1 12 IS *	ACKEY W J ET AL: "CHEMICAL VAPOR POSITION OF OXIDATION RESISTANT HFB2 C COMPOSITE COATINGS" RAMIC ENGINEERING AND SCIENCE COCEEDINGS, COLUMBUS, US, 1. 9, no. 9/10, September 1988 (1988-09-01), pages 23-1231, XP000036430 SN: 0196-6219 abstract * page 1223, line 1 - line 2 * page 1226, chapter "Conclusions" -/	ا 'ممشدا	CO4B C23C
	present search report has been drawn up for all claims		
	ICH Date of completion of the search	ih I	Examiner

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 A : technological background
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 P : intermediate document

SINSPOCID: KEP 124127841 Lis

D : document cited in the application L : document cited for other reasons

8 : member of the same patent family, corresponding document



EUROPEAN SEARCH REPORT

Application Number

EP 02 42 5143

	DOCUMENTS CONSID	OLACOMO ANALISMO			
ategory	Citation of document with of relevant pas	Indication, where appropriate, sages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CI.7)	
	US 5 750 450 A (KAI 12 May 1998 (1998- * abstract * * column 1, line 10 * figure 1 *	JFMAN LARRY ET AL) D5-12)	1,3-6, 15,16		
				TECHNICAL FIELDS SEARCHED (Int.Cl.7)	
	The present search report has	been drawn up for all claims			
	Pace of search	Date of completion of the search		Examiner	
	MUNICH	15 July 2002		Rosenberger, J	
X : partic Y : partic docu A : techr O : non-	ATEGORY OF CITED DOCUMENTS cularly relevant if taken alone cularly relevant if combined with anot ment of the same category nological background written disclosure modifate document	I : theory or prin E : earlier patent after the filing her D : document cit	ciple underlying the in document, but publis	ivention ned on, or	

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EP 02 42 5143

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on

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15-07-2002

	Patent docume cited in search re		Publication date		Patent fam member(s		Publication date
US	4645716	Α	24-02-1987	NONE	**************************************	···	1
us	6110853	A	29-08-2000	DE WO EP	19612926 9736820 0891296	A1	02-10-1997 09-10-1997 20-01-1999
US 	3617358	A	02-11-1971	AT DE FR GB	304728 1794214 1602527 1198745	Al A	15-12-1972 29-07-1971 21-12-1970 15-07-1970
DE	19714433	A	15-10-1998	DE AU AU WO EP JP NO PL	19714433 729953 7209798 9845497 0973955 2001518977 994881 336074	B2 A A1 A1 T	15-10-1998 15-02-2001 30-10-1998 15-10-1998 26-01-2000 16-10-2001 15-11-1999 05-06-2000
US	5750450	A	12-05-1998	NONE			

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